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# Remarks:

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# (54) Process and apparatus for obtaining bulk monocrystalline gallium-containing nitride

(67) The present invention referst to an ammonobasic method for preparing a gallium-containing nitide deviata, in which gallium-containing rededatock is crystallized on at least one crystallization seed in the presence of an alkall metal-containing component in a supercriticalritogen-containing solvent. The method can provide monocystalline gallium-containing nitride crystals having a very high quality.

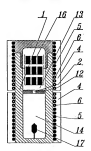


Fig. 9

### Description

[0001] The present invention refers to processes for obtaining a gallium-containing nitride crystal by an ammonobasic method as well as the gallium-containing nitride crystal itself. Furthermore, an apparatus for conducting the various methods is disclosed.

[0002] Optoelectronic devices based on nitrides are usually manufactured on sapphire or allicon carbide substrates that differ from the deposited nitride layers (ex-called heteroepissy). In the most often used Metallo-Organic Chemical Vapor Deposition (MOCVO) method, the deposition of GaN is performed from ammonia and organometallic compounds in the gas phase and the growth rates schieved make it impossible to provide a bulk layer. The application of a buffer layer reduces the dislocation density, but not more than to approx. 10<sup>M</sup>cm<sup>2</sup>, Another method has also been proposed for obtaining but monocrystalling agailtum nitride. This method consists of an epitisal deposition employing halides in a vapor phase and is called Halide Vapor Phase Episaxy (HVPE) ["Optical patterning of GaN films" M.K. Kelly, O. Ambacher, Appl. Phys. Lett. 86 (12) (1998) and "Fabrication of this "Him InGaN Bight-entifing diode membranes" W.S. Wrong, T. Sanda, Appl. Phys. Lett. 86 (10) (1999)]. This method allows for the preparation of GaN substrates having a 2-linch 16 cm id diameter.

[0003] However, their quality is not sufficient for leser diodes, because the dislocation density continues to be approx. 10° to approx. 10°/cm². Recently, the method of Epitaxial Lateral OverGrowth (ELOG) has been used for reducing the dislocation density. In this method the GaN layer is first grown on a sapphire substrate and then a layer with SIO<sub>2</sub> is deposited on it in the form of strips or a lattice. On the thus prepared substrate, in turn, the lateral growth of GaN may be carried out leading to a dislocation density of aporx. 10°/cm².

[0004] The growth of bulk crystals of gaillium nitride and other metals of group XIII (UPAC, 1989) is extremely difficult. Standard methods of crystallization from melt and submetion methods are not applicable because of the decomposition of the nitrides into metals and N<sub>2</sub>. In the High Nitrogen Pressure (HNP) method [\*Prespects for high-pressure crystal growth of III/V nitrides\* S. Porowski et al., Inst. Phys. Conf. Series, 137, 398 (1989) this decorposition is inhibited by the use of nitrogen under the high pressure. They sown of crystals is carried out in molten gaillum, i.e. in the liquid phase, resulting in the production of GaN platelets about 10 mm in size. Sufficient solubility of nitrogen in gaillum requires temperatures of about 1500 °C and introgen pressures in the order of 15 fabor.

[0005]. The use of supercritical ammonia has been proposed to lower the temperature and decrease the pressure during the growth process of intrides. Peters has described the ammonthermal synthesis of aluminium intride (J. Cryst. Growth 104, 411-418 (1990)]. R. Dwillinski et al. have shown, in particular, that it is possible to obtain a fine-crystalline gallium Intride by a synthesis from gallium and runnonia, provided that the latter contains alkali interal amides (CNH-5 or LINH-6). The processes were conducted at temperatures of up to 550° C and under a pressure of 5 kbar, yielding crystalis about 5 µm in size [PAMINON Omethod of BM, NR, and GaM synthesis and crystal growth, \*Proc. EGM-9, Warsew, June 22-24, 1998, MRS Internet Journal of Nitride Semisondudor Research, http://nsr.mij.mrx.org/3/25]. Another supercritical ammonia without, where a fine-crystalline GaM is used as a feedbook topether with a mineralizer consisting of an amide (KNH-5) and a halide (K) also provided for recrystallization or gallium nitride (\*Crystall growth of gallium nitride in supercritical ammonia" JW. Kolis et al., J. Cryst. Growth 222, 431-434 (2001)]. The recrystallization process conducted at 400°C and 3.4 kbar resulted in GaM crystals about 0.5 mm in size. A similar method has also been described in Mat. Res. Sco. Symp. Proc. VI. 445, 367-372 (1988) by JW. Kolis et al. However, using these supercritical ammonia processes, no production of bulk monocrystalline was achieved because no chemical transport processes were observed in the suspercritical solution, in particular no growth on seeds was conducted.

[0006] Therefore, there was a need for an improved method of preparing a gallium-containing nitride crystal.

[0007] The lifetime of optical semiconductor devices depends primarily on the crystalline quality of the optically active

layers, and especially on the surface dislocation density. In case of GaN based laser diddes, it is beneficial to lower the dislocation density in the GaN substrate layer to less than 10%cm², and this has been extremely difficult to achieve using the methods known so far. Therefore, there was a need for gallium-containing nitride crystals having a quality suitable for use as substrates for optoelectronics.

[0008] The subject matter of the present invention is recited in the appended claims. In particular, in one embodiment the present invention refers to a process for obtaining a gallium-containing nitride crystal, comprising the steps of:

 (i) providing a gallium-containing feedstock, an alkali metal-containing component, at least one crystallization seed and a nitrogen-containing solvent in at least one container;

(ii) bringing the nitrogen-containing solvent into a supercritical state;

(iii) at least partially dissolving the gallium-containing feedstock at a first temperature and at a first pressure; and (iv) crystallizing gallium-containing nitride on the crystallization seed at a second temperature and at a second pressure while the nitrogen-containing solvent is in the supercritical state;

wherein at least one of the following criteria is fulfilled:

- (a) the second temperature is higher than the first temperature; and
- (b) the second pressure is lower than the first pressure.
- [0009] In a second embodiment a process for preparing a gallium-containing nitride crystal is described which comprises the steps of:
  - (i) providing a gallium-containing feedstock comprising at least two different components, an alkali metal-containing component, at least one crystallization seed and a nitrogen-containing solvent in a container having a dissolution zone and a crystallization zone, whereby the gallium-containing feedstock is provided in the dissolution zone and the at least one crystallization seed is provided in the crystallization zone;
  - (ii) subsequently bringing the nitrogen-containing solvent into a supercritical state;
  - (iii) subsequently partially dissolving the gallium-containing feedstock at a dissolution temperature and at a dissolution pressure in the dissolution zone, whereby a first component of the gallium-containing feedstock is substantially completely dissolved and a second component of the gallium-containing feedstock as well as the crystallization seed remain substantially undissolved so that an undersaturated or saturated solution with respect to galliumcontaining nitride is obtained;
  - (iv) subsequently setting the conditions in the crystallization zone at a second temperature and at a second pressure so that over-saturation with respect to gallium-containing nitride is obtained and crystallization of gallium-containing nitride occurs on the at least one crystallization seed and simultaneously setting the conditions in the dissolution zone at a first temperature and at a first pressure so that the second component of the gallium-containing feedstock is dissolved:

wherein the second temperature is higher than the first temperature.

- [0010] A gallium-containing nitride crystal obtainable by one of these processes is also described. Further subject matter of the invention are a gallium-containing nitride crystal having a surface area of more than 2 cm2 and having a dislocation density of less than 106 / cm2 and a gallium-containing nitride crystal having a thickness of at least 200 µm and a full width at half maximum (FWHM) of X-ray rocking curve from (0002) plane of 50 arcsec or less.
- [0011] The invention also provides an apparatus for obtaining a gallium-containing nitride crystal comprising an autoclave 1 having an internal space and comprising at least one device 4, 5 for heating the autoclave to at least two zones having different temperatures, wherein the autoclave comprises a device which separates the internal space into a dissolution zone 13 and a crystallization zone 14.
  - [0012] In a yet another embodiment, a process for preparing a bulk monocrystalline gallium-containing nitride in an autoclave is disclosed, which comprises the steps of providing a supercritical ammonia solution containing galliumcontaining nitride with ions of alkali metals, and recrystallizing said gallium-containing nitride selectively on a crystallization seed from said supercritical ammonia solution by means of the negative temperature coefficient of solubility and/or by means of the positive pressure coefficient of solubility.
  - [0013] A process for controlling recrystallization of a gallium-containing nitride in a supercritical ammonia solution which comprises steps of providing a supercritical ammonla solution containing a gallium-containing nitride as a gallium complex with ions of alkali metal and NH3 solvent in an autoclave and decreasing the solubility of said gallium-containing nitride in the supercritical ammonia solution at a temperature less than that of dissolving gallium-containing nitride crystal
- and/or at a pressure higher than that of dissolving gallium-containing nitride crystal is also disclosed. [0014] Fig. 1 shows the dependency of the solubility of gallium-containing nitride in supercritical ammonia that contains potassium amide (with KNH<sub>2</sub>:NH<sub>3</sub>=0.07) on pressure at T=400°C and T=500°C.
- [0015] Fig. 2 shows the diagram of time variations of temperature in an autoclave at constant pressure for Example 1.
- [0016] Fig. 3 shows the diagram of time variations of pressure in an autoclave at constant temperature for Example 2.
- [0017] Fig. 4 shows the diagram of time variations of temperature in an autoclave at constant volume for Example 3.
- [0018] Fig. 5 shows the diagram of time variations of temperature in an autoclave for Example 4. [0019] Fig. 6 shows the diagram of time variations of temperature in an autoclave for Example 5.
- [0020] Fig. 7 shows the diagram of time variations of temperature in an autoclave for Example 6.
- [0021] Fig. 8 shows the diagram of time variations of temperature in an autoclave for Example 7.
- [0022] Fig. 9 shows a schematic axial cross section of an autoclave as employed in many of the examples, mounted in the fumace.
  - [0023] Fig. 10 is a schematic perspective drawing of an apparatus according to the present invention.
  - [0024] Fig. 11 shows the diagram of time variations of temperature in an autoclave at constant volume for Example 8.
- [0025] Fig. 12 shows the diagram of time variations of temperature in an autoclave at constant volume for Example 9.
  - [0026] Fig. 13 shows the diagram of time variations of temperature in an autoclave at constant volume for Example 10. [0027] Fig. 14 shows the diagram of time variations of temperature in an autoclave at constant volume for Examples

11 and 12.

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[0028] Fig. 15 illustrates the postulated theory of the invention.

[0029] In the present invention the following definitions apply:

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Gallium-containing nitride means a nitride of gallium and optonally other element(s) of group XIII (according to tu UPAC, 1989.) In finduces, but is not restricted to, the binary compound Galt, temany compounds such as AIGAN, InGAN and also AlinGAN (The mentioned formulas are only intended to give the components of the nitrides. It is not intended to indicate their relative amounts.

Bulk monocrystalline gallium-containing nitride means a monocrystalline substrate made of gallium-containing nitride means a monocrystalline substrate made of gallium-containing nitride from which e.g. optoelectronic devices such as LED or LD can be formed by epitaxial methods such as MOCVD and HVPE.

Supercritical solvent means a fluid in a supercritical state, it can also contain other components in addition to the solvent itself as long as these components do not substantially influence or disturb the function of the supercritical solvent. In particular, the solvent can contain inso of alkaji metals.

Supercritical solution is used when referring to the supercritical solvent when it contains gallium in a soluble form originating from the dissolution of gallium-containing feedstock.

Dissolution of gallium-containing feedstock means a process (either reversible or irreversible) in which said feedstock is taken up into the supercritical solvent as gallium in a soluble form, possibly as gallium complex compounds.

Gallium complex compounds are complex compounds, in which a gallium atom is a coordination center surrounded by ligands, such as NH<sub>3</sub> molecules or its derivatives, like NH<sub>2</sub>, NH<sup>2</sup>-, etc.

Negative temperature coefficient of solubility means that the solubility of a respective compound is a monotoncially decreasing function of temperature if all other parameters are kept constant. Smillarly, positive pressure coefficient of solubility means that, if all other parameters are kept constant, the solubility is a monotonically increasing function of pressure. In our research we showed that the solubility of gallium-containing intitie in supercritical introgen-containing solvents, such as ammonia, possesses a negative temperature coefficient and a positive pressure coefficient in temperature ranging at least from 300 to 800°C and pressures from 1 to 5.5 kinds.

Over-saturation of supercritical solution with respect to galillum-containing nitride means that the concertration of galium in a soluble form in said solution is higher than that in equilibrum (i.e. it is higher than solubility). In the case of dissolution of galilum-containing nitride in a closed system, such an over-saturation can be achieved by either increasing the temperature and/or decreasing the pressure.

Spontaneous crystallization means an undesired process where nucleation and growth of the gallium-containing nitride from over-saturated supercritical solution take place at any site within an autociave except at the surface of a crystallization seed where the growth is desired. Spontaneous crystallization also comprises nucleation and disoriented growth on the surface of crystallization seed.

Selective crystallization on a seed means a process of crystallization on a seed carried out without spontaneous crystallization.

Autoclave means a closed container which has a reaction chamber where the ammonobasic process according to the present invention is carried out.

0 [0030] The present invention can provide a gallium-containing nitride monocrystal having a large size and a high quality, south gallium-containing nitride crystals can have a surface area of more than 2 cm<sup>2</sup> and a dislocation density of less than 10<sup>4</sup> / cm<sup>2</sup>. Gallium-containing nitride crystals having a thickness of at least 200 µm (preferably at least 500 µm) and a F-WHM of 50 arceece or less can also be obtained. Depending on the crystalization conditions, t possible to obtain gallium-containing nitride crystals having a volume of more than 0.05 cm<sup>3</sup>, preferably more than 0.1 cm<sup>3</sup> using 5 the processes of the invention.

[0031] As was explained above, the gallium-containing nitride crystal is a crystal of nitride of gallium and optionally other element(s) of Group XIII (the numbering of the groups is given according to the IUPAC convention of 1989 throughout this application). These compounds can be represented by the formula AL,Ga<sub>1-x-1</sub>n,N, wherein 05x<1,05y<1,05y<1.

05x+y<1 (preferably 05x<0.5 and 05y<0.5). Although in a preferred embodiment, the gallium-containing nitride is gallium nitride, in a further preferred embodiment part (e.g. up to 50 mol.-%) of the gallium atoms can be replaced by one or more other elements of Groux JMI (sepocially 14 and/or In).

[0032] The gallium-containing nitride may additionally include at least one donor and/or at least one acceptor and/or at least one magnetic dopent a, to a their the optical, electrical and magnetic properties of the substrate. Donor dopents, acceptor dopents and magnetic dopents are well-known in the art and can be selected according to the desired properties of the substrate. Preferably the donor dopents are selected from the group consisting of Si and O. As acceptor donors Mg and Z1 are preferred. Any known magnetic dopant can be included into the substrates of the present invention. A preferred magnetic dopant is Mn and possibly also Ni and Cr. The concentrations of the dopants are well-known in the art and depend on the desired and application of the initiod. Typically the concentrations of these dopants range from 1017 to 1021 / cm², instead of adding dopants as part of the feedstock into the autoclave, dopants can also be included into the gallium-containing nitride crystal from trace amounts of the autoclave material which dissolve during the process of the invention. For example, if the autoclave comprises a nickel alloy then nickel can be included into the gallium-containing nitride crystal from trace amounts of the autoclave material which dissolve during the process of the invention. For example, if the autoclave comprises a nickel alloy then nickel can be included into the gallium-containing nitride crystal.

10033 Due to the preparation process the gallium-containing nitride crystal can also contain alkall elements, usually in an amount of more than about 0.1 ppm. Generally it is desired to keep the alkall elements content tower than 10 ppm, although it is difficult to specify what concentration of alkall metals in gallium-containing nitride has a disadvantageous influence on its prometries.

[0334] It is also possible that halogens are present in the gallium-containing nitride. The halogens can be introduced either intentionally (see a component of the mineralizer) or unintentionally (from inpurities of the mineralizer or the feedstock). It is usually desired to keep the halogen content of the gallium-containing nitride crystal in the range of about

10055] The process of the invention is a supercritical crystallization process, which includes at least two steps: a dissolution step at a first temperature and at a first pressure and a crystallization step at a second repressure. Since generally high pressures and/or high temperatures are involved, the process according to the invention is preferably conducted in an autoclave. The two steps (i.e. the dissolution steps and the crystallization step) can either be conducted separately or can be conducted at least partially simultaneously in the same reaction.

[0038] For conducting the two steps separately, the process can be conducted in one single reactor but the dissolution step is conducted before the crystallization step. In this embodiment the reactor can have the conventional construction of a single chamber. The process of the invention in the two-step embodiment can be conducted using constant pressure and two different temperatures or using constant repressure and two different temperatures. The exact values of pressure and temperature should be selected depending on the feedstock, the specific intrice to be prepared and the solvent. Generally the pressure is into the range of 100 °C to 800 °C, preferably 300 °C to 800 °C, more preferably 400 °C to 550 °C. It two different pressures are employed, the difference in pressure should be from 0.1 bits or both, preferably 400 °C to 500 °C, more preferably 400 °C to 500 °C, introduced the control of the control

(9037) In a preferred embodiment, the dissolution step and the crystallization step are conducted at least partially go (9037) in a preferred pref

50 [0038] A possible construction of a preferred container is given in Fig. 9. For conciseness and ease of understanding in the following, the process will be explained particularly with respect to this preferred embodiment. However, the invention can be conducted with different container constructions as long as the principles outlined in the specification and the claims are adhered to.

[0039] In a preferred embodiment of the invention, the process can be conducted in an apparetus comprising an autoclave 1 having an Internal space and comprising at least on edivice 4, 5 for healing the autoclave 1 or least two zones having different temperatures, wherein the autoclave comprises a device which separates the internal space into a dissolution zone 13 and a crystalization zone 14 (fiveninater size neferred to a "separating device" or "installation"). These two zones having different temperatures should preferably coincide with the dissolution zone 13 and the crystalization zone 14. The device which separates the internal space of the autoclave can be, for example, at least one better size to the space of the space of the success of the space and the control on of the reactor and can be asky determined by a person skilled in the strip.

[0040] In none embodiment, two different hearing devices can be employed, the position of which preferably corresponds to the dissolution zone 13 and the reputalization zone 14 theower, it has been observed that transport of gallumin in a soluble form from the dissolution zone 13 to the crystalization zone 14 can be further improved if a cooling means 6 is present between the first and the second hearing devices and is located at approximately the position of the separating device. The cooling means 6 can be realized by liquid (e.g., water) cooling or preferably by fan cooling. The healing devices can be powered electrically, by either indevide or, preferably, by resistive healing means. Use of a heating-cooling, heating configuration gives wider possibilities informing the desired temperature distribution within the autoclave. For example, it enables to obtain a low temperature gradient with in most of the crystalization zone 14 and a low temperature gradient within most of the region of the belff e12.

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(O41) When the process of the present invention is conducted a gallium-containing feedstock, an alkali metal-containing component, at least one crystallization seed and a introgen-containing solvent are provided in at least one container. In the preferred apparatus described above, the gallium-containing feedstock it is placed in the dissolution zone 13 and the at least one crystallization seed 17 is placed in the crystallization zone 14. The alkali metal-containing component is also preferably placed in the dissolution zone. Then the nitrogen-containing solvent is added to the container, which is then closed. Subsequently the nitrogen-containing solvent is brought into a supercritical state, e.g. by increasing pressure and/or heat.

[0042] In the present invention any materials containing gallium, which are soluble in the superritical solvent under the conditions of the present invention, can be used as a spallium containing feedstock. Typically the gallium-containing interestors, Propietally the gallium-containing interestors, Propietally the gallium-containing interestors, Propietally the gallium-containing interestors and inter-drealized comproused, by drieds, andies, inclose, andied-indee, such as called-gallium-containing letters and inter-drealized comproused, by drieds, and interest gallium-nitride GaN, azides such as Ga(N+)<sub>2</sub>, imides such as Ga(N+)<sub>2</sub>, by thirdses us das Ga(N+)<sub>2</sub>, by thirdses under the gallium-containing allium, metallic gallium-drealized and mixtures thereof. Most preferably, the feedstock is metallic gallium and gallium intride. It elements of Group XIII other than gallium and be present in the gallium-containing allium, containing allium-containing al

[0043] The form of the feedstock is not particularly limited and it can be in the form of one or more pieces or in the form of a powder. If the feedstock is in the form of a powder, care should be taken that individual powder particles are not transported from the dissolution zone to the crystallization zone, where they can cause irregular crystallization. It is preferable that the feedstock is in one or more pieces and that the surface area of the feedstock is larger than that of the crystallization seed.

[0044] The nitrogen-containing solvent employed in the present invention must be able to form a supercritical fluid, in which gaillum to be dissolved in the presence of alkali metal ions. Preferably the solvent is ammonia, a derivative theories of mixtures thereof. An example of a suitable ammonia derivative is hydrazine. Most preferably the solvent is ammonia. To reduce corrosion of the reactor and to avoid side-reactions, habgens e.g., in the form of haldes are preferably not intentionally added into the container. Although traces of habgens may be introduced into the system in the form of unavoidable impurities of the starting materials, care should be taken to keep the amount of halogen as low as possible. Due to the use of a nitrogen-containing solvent souch as ammonia it is not necessary to include nitride compounds into the feedstock. Metallic gaillum (or aluminium or indium) can be employed as the feedstock while the solvent provides the nitrogen required for the nitride formation.

[0045] It has been observed that the solubility of gallium-containing feedstock, such as gailium and corresponding elements of Group XIII and/or their compounds, can be significantly improved by the presence of at least one type of alkali metal-containing component as a solubilization aid ("mineralizer"). Lithium, sodium and potassium are preferred as alkali metals, wherein sodium and potassium are more preferred. The mineralizer can be added to the supercritical solvent in elemental form or preferably in the form of as compound (such as a sail). Generally the choice of the mineralizer depends on the solvent employed in the process. According to our investigations, alkali metal having a smaller ion radius can provide lower solubility of gallium-containing intide in the supercritical solvent than that obtained with alkali metals having a larger ion radius. For example, if the mineralizer is in the form of a compound such as a sait, it is preferably in the form of an alkali metal families such as MM, as alkali metal minde such as MM, as a make in metal families such as MM, as a make in metal families out as a MM, and a selected so as a to ansure adequate levels of solubility of both feedstock (the starting material) and gailium-containing nitride (the resulting product). It is usually in the range of 1: 200 to 1: 2, in the terms of the most of the metal ion based on the most of the metal ion bas

[0046] The presence of the alkali metal ions in the process can lead to alkali metal in the thus prepared substrates.

It is possible that the amount of alkali metal is more than about 0.1 ppm, even more than 10 ppm. However, in these amounts the alkali metals do not detrimentally effect the properties of the substrates. It has been found that even at an alkali metal content of 500 ppm, the operational parameters of the substrate according to the invention are still satisfactory. [0047] The dissolved feedstock crystallizes in the crystallization step under the low solubility conditions on the crystallization step under the low solubility conditions on the crystallization step under the low solubility conditions on the crystallization step under the low solubility conditions on the crystallization step under the low solubility conditions on the crystallization step under the formation of stoichiometric gallium-containing nitride in the form of a bulk monocystalline layer on the crystallization seed(e) and in particular leads to the formation of stoichiometric gallium-containing nitride in the form of a bulk monocystalline layer on the crystallization seed(e).

[0048] Various crystals can be used as crystallization seeds in the present invention, however, it is preferred that chemical and crystallicographic constitution of the crystallization seeds is similar to those of the desired layer of bulk monocrystalline gallium-containing nitride. The relational profit of the dissolved preferrably compress a crystalline layer of gallium-containing nitride. To facilitate crystallization of the dissolved redeatock, the discontained send that crystallization gallium-containing nitride. To facilitate crystallization of the dissolved redeatock, the discontained send that crystallization seed gardents, the discontained send to the crystalline layer of or more and thickness of 100 um or part of the dissolved for the discontained to the crystalline layer.

[0049] After the starting materials have been introduced into the container and the nitrogen-containing solvent has been brought into its supercritical state, the gallium-containing feadatock is at least partially dissolved at a first temperature and a first pressure, a.g. in the dissolution zone of an autoclave. Gallium-containing initide crystallization seed (e.g. in the crystallization zone of an autoclave) at a second temperature and at a second pressure while the nitrogen-containing solvent is in the supercritical state, wherein the second itemperature is higher than the first temperature and/or the second pressure is lower than the first pressure. If the dissolution and the crystallization steps take place similar abouts in the same container, the second pressure is essentially equal to the first pressure.

(1950) This is possible since the solubility of gallium-containing nitride under the conditions of the present invention shows a negative temperature coefficient and a positive pressure coefficient in the presence of affair metal lors. Without withing to be bound by theory, it is postulated that the following processes occur. In the dissolution 20ne, the temperature and pressure are selected such that the gallium-containing feededstock is dissolved and the introper-containing solution is undersaturated with respect to gallium-containing introde. In the crystallization zone, the temperature and pressure are selected such that the solution, atthough it contains approximately the seme concentration of gallium as in the dissolution zone, is over-esturated with respect to gallium-containing nitride. Therefore, crystallization of gallium containing intride on the crystallization gallium-containing nitride on the crystallization gallium-containing printed expected in the containing the seminated of the crystallization gallium-containing the investment of the crystallization gallium-containing intride on the crystallization gallium-containing intride interest of the crystallization ground crystallization gallium-containing intride interest of the crystallization zone. In the present invention this erferred to as the femical transported of gallium-containing intride in the superviolate solution. It is produted that the soluble form of gallium is a gallium complex compound with a Ga atom in the coordination center surrounded by ligands, such as NH<sub>2</sub> molecules or its derivatives or its derivatives, its NH<sub>2</sub>. NH-2.

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[0051] This theory is equally applicable for all galillum-containing nitrides, such as AGAN, InGAN and AIMGAN as well as GaN (The mentioned formulas are only intended to give the components of the nitrides. It is not intended to indicate their relative amounts). In the case of nitrides other than gallium nitride aluminum and/or indum in a soluble form also have to be present in the supervictical solution.

[0052] In a preferred embodiment of the invention, the galilum-containing feedstock is dissolved in at least two steps, in this ambodiment, the galilum-containing feedstock, generally comprises two kinds of starting materials which differ in solubility. The difference in solubility can be achieved chemically (e.g. by selecting two different chemical compounds or physically (e.g. by selecting two fifteents chemical compounds under the compound having for example different surface areas, like micro-cystalline porder and large crystals). In a preferred embodiment, the galilum-containing feedstock comprises two different chemical compounds such as metalic galilum and galilum nitride which dissolve at different rates, in a first dissolution temperature and at a dissolution pressure in the dissolution. The dissolution is understand that the dissolution pressure, which can be set only in the dissolution area or preferably in the whole containing, are selected so that the second component of the galilum-containing feedstock and the crystallization seed(a) remain substantially undissolved. This first dissolution step result is an undersaturated of art most asturated solution (preferably undersaturated solution) from 150 °C to 350 °C, preferably from 150 °C to 350 °C. The dissolution pressure can be ol. 1 kbar to 8 kbar, preferably from 1, 1 kbar to 8 kbar, preferably from 1, 1 kbar to 8 kbar.

(0033) Subsequently the conditions in the crystallization zone are set at a second prepretary and a conservation of the conservation of the crystallization can be a conservation of the crystallization can be a conservation of the crystallization of the

and the crystallization step, the system should be in a stationary state so that the concentration of gallium in the supercritical solution remains substantially constant, i.e. sproxymately the same amount of gallium should be discoved per unit of time as is crystallized in the same such of time. This allows for the growth of gallium-containing nitride crystals of sealing in calling the contraction of the c

[0054] Typical pressures for the crystallization step and the second dissolution step are in the range of 1 to 10 kbar, preferably 1 to 5.5 kbar and more perferably 1.5 to 8 kbar. The temperature is pomerally in the range of 10 to 800°C, preferably 300 to 800°C, more preferably 400 to 550°C. The difference in temperature should be at least 1°C, and is preferably from 5°C to 150°C. As explained above, the temperature difference between the dissolution zone and crystallization zone should be controlled so as to ensure chemical transport in the supercritical solution, which takes

place through convection.

[0055] In the process of the invention, the crystallization should take place selectively on the crystallization seed and not on a wall of the container. Therefore, the over-saturation extent with respect to the galilum-containing nitride in the supercritical solution in the crystallization zone should be controlled on as to be below the spontaneous crystallization level where crystallization takes place on a wall of the autoclave and/or disordered growth occurs on the seed, i.e. the level at which poptonaneous crystallization cours. This can be achieved by adjusting the chemical transport rate and/or the crystallization temperature and/or crystallization pressure. The otherwise the provided the properties of the convective flow from the dissolution zone to the crystallization zone, which can be controlled bythe temperature difference between the dissolution zone and the crystallization zone, the size of the opening(s) of baffle(s) between the dissolution zone and the crystallization zone, the size of the opening(s) of baffle(s) between the dissolution zone and the crystallization zone, the size of the opening(s) of baffle(s) between the dissolution zone and the crystallization zone, the size of the opening(s) of baffle(s) between the dissolution zone and the crystallization zone.

20 [0056] The performed tests showed that the best bulk monocrystalline gallium nitride obtained had a dislocation density close to 10<sup>4</sup>0cm<sup>2</sup> and simultaneously a FWHM of X-ray rocking curve from (0002) plane below 60 arcsec. These crystals possess an appropriate quality and durability for optical semiconductor devices. The gallium-containing nitride of the present invention typically has a wurzite structure.

[0057] Feedstock material for use in the present invention can also be prepared using a method similar to those described above. The method involves the steps of:

- (i) providing a gallium-containing feedstock, an alkali metal-containing component, at least one crystallization seed and a nitrogen-containing solvent in a container having at least one zone;
- (ii) subsequently bringing the nitrogen-containing solvent into a supercritical state;
- (ii) subsequently dissolving the gallium-containing feedstock (such as metallic gallium or aluminium or indium, perferably metallic gallium) at aliciasolution pressure, whereby the gallium-containing feedstock is substantially completely dissolved and the crystallization seed remains substantially undissolved so that an undersaturated solution with respect to callium-containing initiation inhibits in obtained.
- (iv) subsequently setting the conditions in at least part of the container at a second temperature and at a second pressure so that over-saturation with respect to galitum-containing nitride is obtained and crystallization of galliumcontaining nitride occurs on the at least one crystallization seed;

wherein the second temperature is higher than the dissolution temperature.

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[0058] In this embodiment the comments given above with respect to the individual components, process parameters, to etc. also apply. Freferably during the crystalization step in this embodiment the conditions in the whole container are set at the second temperature and the second pressure.

[0059] Gallium-containing nitride exhibits good solubility in a supercritical nitrogen-containing solvent (e.g. ammonia), provided alkali metals or their compounds, such as KNH₂, are introduced into it. Fig. 1 shows the solubility of gallium-containing nitride in a supercritical solvent versus pressure for temperatures of 400 and 500°C wherein the solubility is defined by the moiar percentage: S<sub>m</sub>= GaNevent® (KNH₂, NH₃) 100%. In the present case the solvent is supercritical ammonia containing KNH₂ in a molar ratio x KNH₂, NH₃ equal to 0.07. For this case S<sub>m</sub> should be a smooth function of only three parameters: temperature, pressure, and molar ratio of mineralizer (i.e. S<sub>m</sub> = S<sub>m</sub>(T, p, x)). Small changes of S<sub>m</sub> can be expressed as:

$$\Delta S_m \approx (\partial S_m/\partial T)|_{D_X} \Delta T + (\partial S_m/\partial p)|_{T_X} \Delta p + (\partial S_m/\partial x)|_{T_X} \Delta x$$

where the partial differentials (e.g. (∂S<sub>m</sub>(∂T))<sub>p,x</sub>) determine the behavior of S<sub>m</sub> with variation of its parameters (e.g. T). In this specification the partial differentials are called "coefficients" (e.g. (∂S<sub>m</sub>(∂T))<sub>p,x</sub> is a "temperature coefficient" of solubility or "temperature coefficient".

[0060] The diagram shown in Fig. 1 illustrates that the solubility increases with pressure and decreases with temper-

ature, which means that it possesses a negative temperature coefficient and a positive pressure coefficient. Such features allow obtaining a bulk monocrystalize julium-containing nitride by dissolution in the higher solubility conditions, and crystalization in the lower solubility conditions, and crystalization in the lower solubility conditions. In perioduar, the negative temperature coefficient means that, in the presence of a temperature gradient, the chemical transport of gallium in a solubile form can take piace from the dissolution zone having a lower temperature to the crystalization zone having a likely temperature.

zone having a lower temperature to the crystalization 2 one having a night emperature. 
[0601] The process according to invention allows the growth of bulk monocrystalline gallium-containing nitride crystals 
on the crystallization seed and leads in particular to the formation of stoichlometric gallium-containing nitride, obtained 
in the form of a bulk immocrystalline layer grown on a gallium-containing nitride crystallization seed. Shoe such a 
monocrystal is obtained in a supercritical solution that contains ions of alkali metals, it can contain alkali metals in a 
quantity higher than 0.1 ppm. Because it is desired to maintain a purely basic character of the supercritical solution, 
mainly in order to avoid corrosion of the appearatus, halides are preferably not intentionally introduced into the solvent. 
The process of the invention can also provide a bulk monocrystalline gallium-containing nitride crystal in gride crystal may be doped with donor and/or occeptor and/or magnetic departs. These dopants can modify 
optical, electric and magnetic properties of the gallium-containing nitride crystal and the state of the particular 
properties, the bulk monocrystalline gallium-domaining nitride crystal and can have a disclosation density below 10%/cm², 
preferably below 10%/cm², or most preferably below 10%/cm², Desides, the FWHM of the X-ray rocking curve from (0002) 
plane can be below 600 arcsec, preferably below 000 arcsec, and most preferably below 60 arcsec. The best bulk 
monocrystalline gallium nitride obtained may have a dislocation density below 60 arcsec. The best bulk 
monocrystalline gallium office or preferably below 00 arcsec, and most preferably below 60 arcsec. The best bulk 
monocrystalline gallium-domain of the crystal row of arcsec. The best bulk 
monocrystalline gallium office or preferably below 00 arcsec, and most preferably below 60 arcsec. The best bulk 
monocrystalline gallium office or preferably below 00 arcsec, and most preferably below 60 arcsec. The best bulk 
monocrystalline galli

[0062] Due to their good crystalline quality the gallium-containing nitride crystals obtained in the present invention may be used as a substrate material for optoelectronic semiconductor devices based on nitrides, in particular for laser diodes.

[0063] The following examples are intended to illustrate the invention and should not be construed as being limiting.

### **EXAMPLES**

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[0064] The dislocation density can be measured by theso-called EPD method (Etch Pit Density) and subsequent evaluation using a microscope

[0065] The FWHM of the X-ray rocking curve can be determined by X-ray diffraction analysis.

[0056] Since it is not possible to readily measure the temperature in an autoclave while in use under supercritical conditions, the temperature in the autoclave was estimated by the following method. The outside of the autoclave was estimated by the following method. The outside or the autoclave is equipped with thermocouples near the dissolution zone and the crystallization zone. For the calibration, additional temperature is the enterpoly autoclave was then heated stepwise to various temperatures and the values of the temperature of the emperature and the values of the temperature of the experiment is naised the autoclave and outside the autoclave were measured and tabulated. For example, if the temperature of the crystallization zone is determined to be 500. °C and the temperature of the dissolution zone is 400. °C respectively, it is assumed that under supercritical conditions the temperature of the crystallization/dissolution zone is 400. °C will also be 500. °C/400. °C when the representative in the crystallization/dissolution zone was under the crystallization zone. The crystallization zone is 400. °C will also be 500. °C/400. °C when temperature in the crystallization zone is 400. °C will also be 500. °C/400. °C when temperature difference between the two zones can be lower due to effective heat transfer through the supercritical solution.

### Example 1

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[0067] Two crucibles were placed into a high-pressure autodave having a volume of 10.9 cm². The autodave mannaturactured according to a known design [H. Jacoba, D. Schmidt, Current Topis in Miteriale Science, vol. 8, ed. E. Kaidis (North-Holland, Amsterdam, 1881), 381) One of the crucibles contained 0.4 g of gailium nitride in the form of 0.1 munitive place produced by the HVPE method as feedstock, while the other contained a gailium nitride in the form of 0.1 multive places weighing 0.1 g. The seed was also obtained by the HVPE method. Further, 0.72 g of metallic potassium of N purty were placed in the autodave, the autodave was filled with 4.8 g of ammonia and then closed. The autodave was put into a furnace and heated to a temperature of 40°C. The pressure within the autodave was placed in the autodave was maintained under these conditions for another 6 days (Fig. 2). As a result of this process, in which the dissolution of crystallization steps were separated in time, the feedstock was completely dissolved and the recystalization of gailium nidride in the form of a layer took place on the partialty dissolved seed. The two-sided monocrystalline layers had a total thickness of about 4.4 mm.

### Example 2

10058] Two crucibles were put into the above-mentioned high-pressure autoclave having a volume of 10.9 cm<sup>3</sup>. One of the crucibles contained 0.44 g of galilum nitride in the form of 1.mm this plates produced by the HVPE method as feedstock, and the other contained a galilum nitride seed of a double thickness weighing 0.1g, also obtained by the HVPE method. But the third of the contained a galilum nitride upon the produced of the successive weighing 0.1g, also obtained by the HVPE method. But the contained the successive weighing 0.1g, also obtained by the MVB 5.43 g of metallier potential of the successive was put into a furnees and heated to a temperature of 50°C. The pressure within the autoclave was 3.5 kbar. After 2 days the pressure was lowered to 2 kbar, while the temperature was maintained at the 50°C level and the autoclave was maintained under these conditions for another 4 days (Fig. 3). As a result of this process, the feedstock was completely dissolved and the recrystalization of galilum nitride took place on the partially dissolved asod. The two-sided monocrystalline dayses had a total thickness of about 0.25 mm.

### Example 3

[0063] Two crucibies were placed into the above-mentioned high-pressure autoclave having a volume of 10.9 cm², fone of the crucibles contained 0.3 g of the feedstock in the form of metallic gaillaum of Rb purity and the other contained a 0.1 gaillaum nitride seed obtained by the HVPE method. Further, 0.6 g of metallic politics six of Ab purity was placed in the autoclave; the autoclave was filled with 4 g of ammonia and then closed. The autoclave was put into a furnace and heated to a temperature of 20°Co. After 2 days the temperature was increased to 50°Cc, while the pressure was maintained at the 2 kbar level and the autoclave was mentiatised in the seconditions for further 4 days (Fig. 4). As a result of this process, the feedstock was completely dissolved and the crystallization of gaillum nitride took place on the seed. The two-sided monocrystallic levers had a total tribickness of shout 0.3 mm.

# Example 4

[0070] This is an example of a process, in which the dissolution and crystallization steps take place simultaneously (recrystallization process). In this example and all the following an apparatus is used which is schematically shown in Fig. 9 and Fig. 10. The basic unit of the apparatus is the autoclave 1, which in this Example has a volume of 35.6 cm<sup>3</sup>. The autoclave 1 is equipped with an separating device 2 which allows for chemical transport of the solvent in the supercritical solution inside the autoclave 1. For this purpose, the autoclave 1 is put into a chamber 3 of a set of two furnaces 4 provided with heating devices 5 and a cooling device 6. The autoclave 1 is secured in a desired position with respect to the furnaces 4 by means of a screw-type blocking device 7. The furnaces 4 are mounted on a bed 8 and are secured by means of steel tapes 9 wrapped around the furnaces 4 and the bed 8. The bed 8 together with the set of furnaces 4 is rotationally mounted in base 10 and is secured in a desired angular position by means of a pin interlock 11. In the autoclave 1, placed in the set of furnaces 4, the convective flow of supercritical solution takes place as determined by the separating device 2. The separating device 2 is in the form of a horizontal baffle 12 having a circumferential opening. The baffle 12 separates the dissolution zone 13 from the crystallization zone 14 in the autoclave 1, and enables, together with the adjustable tilting angle of the autoclave 1, controlling of speed and type of convective flow. The temperature level of the Individual zones in the autoclave 1 is controlled by means of a control system 15 operating the furnaces 4. In the autoclave 1, the dissolution zone 13 coincides with the low-temperature zone of the set of furnaces 4 and is located above the horizontal baffle 12 and the feedstock 16 is put into this zone 13. On the other hand, the crystallization zone 14 coincides with the high-temperature zone of the set of furnaces 4 and it is located below the horizontal baffle 12. The crystallization seed 17 is mounted in this zone 14. The mounting location of the crystallization seed 17 is below the intersection of the rising and descending convective streams.

45 [0071]. An amount of 3 og of gallium intide produced by the HVPE method was pleaced in the high-pressure autoclave described above, which was set in the horizontal position. This gallium intide had the form of plates of about 0.2 mm thickness, and it was distributed (roughly uniformly) in equal portions in the dissolution zone 13 and the crystallization zone 14. The portion placed in the dissolution zone 13 played the note of redestock, whereas the portion placed in the operation zone 14 played the role of crystallization seeds. Metallic potassium of 4N purity was also added in a quantity of 2-q. Than the autoclave 1 was filled with 15 9g of amomonia (6N), closed, put into a set of transces 4 and heated to a temperature of 450°C. The pressure inside the autoclave 1 was approx. 2 kbar. During this stage, which lasted one day, partial dissolution of gallium intide was carried out in both zones. Then the temperature of the cystallization zone 14 was increased to 500°C while the temperature of the dissolution zone 13 was lowered to 400°C and the autoclave 1 was kept in these conditions for 6 more days (Fig. 5). As a final result of this process, partial dissolution 500 for the feedstock in the dissolution zone 13 and crystallization of gallium nitride on the gallium nitride seeds in the crystallization zone 14 by lines.

# Example 5

[0072] The above-mentioned high pressure autoclave 1 having a volume of 3.5 c cm<sup>3</sup> was charged with feedstock in the form of a 3.0 go pellor of siretering alliam intride (introduced into the dissolution zone 13), two seeds of gallium nitride obtained by the HVPE method and having the form of plates having a thickness of 0.4 mm and a total weight of 0.1g (introduced into the crystallization zone 14), as well as with 2.4g of metallic potassium of 4N purity. Then the autoclave was filled with 1.5g of ammonial 6N) and closed. The autoclave 1 was then put into a set of furnaces 4 and hested to 450°C. The pressure inside the autoclave was about 2 kbar. After an entire day the temperature of the crystallization zone 14 was raised to 480°C, while the temperature of the dissolution zone 13 was lowered to 420°C and the autoclave was maintained under these conditions for 6 more days (see Fig. 6). As a result of the process the feedstock was partially dissolved in the dissolution zone 13 and solventille layers had a for gallium nitride crystallized on the seeds in the crystallization zone 14. The two-sided monocytealline layers had solt his charge to about 0.2 mm.

### Example 6

[0073] The above-mentioned high pressure autoclave 1 having a volume of 3.5 ft cm<sup>2</sup> (see Fig. 9) was changed with 1.6g of feedstock in the form of galilum nitride produced by the HVPE method and having the form of platies having a thickness of about 0.2 mm (introduced into the dissolution zone 13), three galilum-nitride seeds of a thickness of about 0.55 mm and a total weight of 0.6g, also obtained by the HVPE method (introduced into the crystalization zone 14), as well as with 3.56 g of metallic potassistum of 4N punty. The autoclave 1 was tilled with 14.5g of ammonia (6N) and closed. Then the autoclave 1 was put into a set of turnaces 4 and heated to 42°C. The pressure inside the autoclave approx. 1.5 bits. After an entiric day the temperature of the dissolution zone 13 was increased to 45°C and the autoclave was kept in these conditions for 8 more way (see Fig. 7). After the processes, the feedstock was found to be partially dissolved in the dissolution zone 13 and agalilum intride had crystallization those sed of the HVPE GaN in the crystalization zone 14. The two-sided monocrystalline lavers had a total thickness of about 0.15 mm.

### Example 7

[0074] The above-mentioned high pressure autoclave 1 having a volume of 35.6 cm² (see Fig. 9) was charged in its dissolution zone 13 with 2g of feedstock in the form of galillum nitride produced by the HVPE method and having the form of plates having a thickness of about 0.2 mm, and 0.47g of metallic potassium of 44 Purity, and 11 his crystallization zone 14 with three GaN seeds of a thickness of about 0.3 mm and a total weight of about 0.3g also obtained by the HVPE method. The autoclave was filled with 16.5g of ammonia (6.9k) and closed. Then tha autoclave 1 was put into a set of furnaces 4 and heated to 500°C. The pressure inside the autoclave was approx. 3 bkar. After an entire day the temperature in the dissolution zone 14 was reduced to 450°C while the temperature in the crystallization zone 14 was reduced to 450°C while the temperature in the dissolution zone 14 was reduced to 450°C while the temperature in the dissolution zone 14 was reduced to 450°C while the temperature in the dissolution zone 13 ms are dissoluted in the crystallization zone 4 was the crystallization zone 14 will be a condition to the partial day (see Fig. 8). After the process, the feedstock was found to be partially dissolved in the dissolution zone 13 and gallium nitride had crystallization zone 14 ms.

# Example 8

[Q075] An amount of 1.0g of gallium nitride produced by the HVPE method was put into the dissolution zone 13 of the high-pressure autorioave 1 having a thickness of 100 μm and a surface area of 2.5 cm², obtained by the HVPE method, was pleaded. Then the autorioave, are propagation and a surface area of 2.5 cm², obtained by the HVPE method, was pleaded. Then the autorioave was changed with 1.2 g of metallic gallium of 6N purity and 2.2 g of metallic pallium of 6N purity and 2.2 g of metallic pallium and extended to the surface area of 4N purity. Subsequently, the autoclave 1 was filled with 15.9g of ammonia (6N), closed, put into a set of furnaces 4 and 4N purity. Subsequently, the autoclave 1 was filled with 15.9g of ammonia (6N), closed, put into a set of furnaces 4 and extended to a temperature or 200°C. After 3 days - during which period metallic gallium was dissolved in the supercritical solution - the temperature was increased to 50°C while the setuled in a pressure of about 2.3 kbar. The next day, the crystallization pare temperature was increased to 50°C while the temperature of the dissolution zone 13 was lovered to 30°C and the autoclave 1 was kept in these conditions for the next 20 days (see Fig. 11). As a result of this process, the partial dissolution of the material in the dissolution zone 13 and the growth of the gallium mirtide one the gallium mirtide ased in the crystallization zone 14 took place. The resulting crystal of gallium nitride shaving a total thickness of 350 μm was obtained in the form of two-sided monocrystalline leyers.

### Example 9

[0076] An amount of 3.0g of gallium nitride in the form of a sintered gallium nitride pellet was put into the dissolution

zone 13 of high-pressure autoclave 1 having a volume of 35.6 cm² (see Fig. 9). In the crystallization zone 14 of the autoclave, a crystallization zone 14 of the autoclave, a crystallization zone 14 of the autoclave, a crystallization zone 14 of the surface area of 22 cm² was placed. Then the autoclave was charged with 2.3 g of metallic potassium of 4k purity. Subsequently, the autoclave 1 was filled with 15.9g of ammonia (8k), closed, put into a set of turnaces 4 and heated to a temperature of 25°C in order to partially dissolve the sintered Gan Pielet and obtain a preliminary saturation of a supercritical solution with gallium in a soluble form. After two days, the temperature of the crystallization zone 14 was increased to 50°C while the temperature of the dissolution zone 13 was lovered to 42°C and the autoclave 1 was kept in these conditions for the next 20 days (see Fig. 12). As a result of this process, partial dissolution or the material in the dissolution zone 13 and growth of gallium intirde on the gallium intiride sed took place in the crystallization zone 14. A crystal of gallium nitride having a total thickness of 500 μm was obtained in the form of two-sided monocrystalline

### Example 10

[0077]. An amount of 0.5g of gellium näride plates having an average thickness of about 120 µm, produced by the HVPE method, were put into the dissolution zone 13 of high-pressure autocaives 1 waiving a volume of 35.6 cm³. In the crystallization zone 14 of the autoclave, three crystallization seeds of gallium nitride obtained by the HVPE method were placed. The crystallization seeds had thickness of about 120 µm and a total surface area of 1.5 cm². Then the autoclave was charged with 0.41 go' metallic limitum of 31 punit. Subsequently, the autoclave 1 was filled with 14.4g of ammonia (5N), closed, put into a set of rumaces 4 and heated so that the temperature of the crystallization zone 14 was increased to 550°C and the temperature of the dissolution zone 0 and 13 was increased to 450°C. The resulting pressure was about 25 kbar. The autoclave 1 was kept in these conditions for the next 8 days (see Fig. 13). As a result of this process, partial dissolution of the material in the dissolution zone 13 and growth of gallium nitride on the gallium nitride seeds in the crystallization zone 14 took place. The resulting crystals of gallium nitride had a thickness of 40 µm and were in the form of two-deided monocrystalline layers.

### Example 11

[0078] An amount of 0.5g of gallium nitride having an average thickness of about 120 µm, produced by the HVPE method, was placed into the dissolution zone 13 of high-pressure autociave 1 having a volume of 3.5 a cm<sup>3</sup>. In the crystallization seeds of gallium nitride obtained by the HVPE method were placed. The crystallization seeds had a thickness of 120 µm and a total surface area of 1.5 cm<sup>2</sup>. Then the autociave was charged with .071 g of methalic gallium of 8 Pb purity, 5 busequently, the autociave 1 was filled with 14.5g of ammonia (5N), closed, put into a set of furnaces 4 and heated to a temperature of 20°C. After 1 day - during which period methalic gallium was dissolved in the supportificial solution 1 he autociave 1 was heated so that the temperature in the cissolution zone was increased to 50°C, while the temperature in the dissolution zone was increased to 50°C, while the temperature in the dissolution zone was increased to 50°C, while the temperature in the dissolution zone the next 8 days (see Fig. 14). As a result of this process, partial dissolution of the material in the dissolution zone 13 and growth of gallium mitride on the gallium nitride seeds in the crystallization zone 14 took place. The resulting crystals of gallium nitride was challed in the crystallization zone 14 took place. The resulting crystals of gallium nitride on the gallium nitride on five order on five added to the process partial dissolution zone 14 took place. The resulting crystals of gallium nitride on the gallium nitride on five order having a total thickness of 400 µm.

# Example 12

[0079] An amount of 0.5g of gallium nitride having an average thickness of about 120 μm, produced by the HVPE of the thind, was placed into the dissolution zone 13 of the sligh-pressure autoclave 1 having a volume of 3.5, cm³. In the crystallization zone 14 of the autoclave, three crystallization seeds of pallium intrice obtained by the HVPE method were placed. The crystallization seeds had a thickness of 120 μm and a total surface area of 1.5 cm². Then the autoclave was charged with 0.2g of gallium amide and 1.4g of metallic soldium of 3N purity. Subsequently, the autoclave 1 was filled with 1.4 6g of ammonia (SN), closed, put into a set of furnaces 4 and heated to a temperature of 20°C. After 1 day of the control of the subsequently the autoclave 1 was bested so that the temperature in the crystallization zone was increased to 40°C. The resulting pressure was about 2.3 kbar. The autoclave 1 was kept in these conditions for the next 8 days (see also Fig. 1.4). As a result of the process, partial dissolution of the material in the dissolution zone 13 and growth of gallium nitride over in the form of two-sided monocrystalliar layers having a total thickness of 49 μm.

### Example 13

10080] One crucible was placed into the above-mentioned high-pressure autoclave having a volume of 10.9 cm². The crucible contained 0.3 g of the feededock in the form of metallic galitim of 8D puirty. Most three galiturnitide seeds having a thickness of about 0.5mm and a total mass of 0.2g, all obtained by the HVPE method, were suspended within the autoclave. Further, 0.5 g of memoral. The autoclave was put into a funnece and heated to a temperature of 200°C, where the pressure was about 2.5 kbar. After 1 day the temperature was increased to 500°C, while the pressure increased up to 500°C, where the pressure increased up to 500°C, where the pressure increased up to 500°C, while the pressure increased up to 500°C, where the pressure increased up to 500°C, while the pressure in

ray rocking curve from the (0002) plane at the gallium-terminated side was 43 arcsec, while at the nitrogen-terminated side it was 927 arcsec.

[0081] The monocrystalline gallium nitride layers have a wurzite structure like in all of the other examples.

# Claims

- 1. A process for obtaining a bulk monocrystalline gallium-containing nitride crystal, wherein it is performed in an autoclave, in the environment of a supervitical solvent containing ions of a kall metals, wherein a gallium-containing interfedence disolved in said supervitical solvent to form a superoritical solution, and the gallium-containing nitride becomes crystallized from the supercritical solution on the surface of a crystallization seed at a temperature higher and/or pressure lower than that of the feedstock dissolution in the supercritical solvent.
- The process according to claim 1, wherein said process comprises the steps of dissolving the gallium-containing feedstock and a separate step of transferring the supercritical solution to the higher temperature and/or to the lower pressure.
- 3. The process according to claim 1 wherein said process comprises the step of simultaneous creation of at least two zones of different temperatures, said gaillum-containing feedstock is placed in the dissolution zone of the lower temperature, while the drystallization seed is placed in the crystallization zone of the higher temperature.
  - The process according to claim 3, wherein said temperature difference between said dissolution zone and said crystallization zone is controlled so as to ensure chemical transport in the supercritical solution.
    - The process according to claim 4, wherein said chemical transport in the supercritical solution takes place through convection in the autoclave.
- The process according to claim 4, wherein said temperature difference between the dissolution zone and the crystallization zone is greater than 1°C.
  - The process according to claim 1, wherein said gallium-containing nitride crystal has the formula Al<sub>x</sub>Ga<sub>1-x-y</sub>in<sub>y</sub>N, where 0≤x<1, 0≤y<1, 0≤x+y<1.</li>
  - The process according to claim 1, wherein said gallium-containing nitride crystal contains dopants of a donor and/or acceptor and/or magnetic type.
  - The process according to claim 1, wherein said supercritical solvent contains NH<sub>3</sub> and/or its derivatives.
  - 10. The process' according to claim 1, wherein said supercritical solvent contains sodium and/or potassium ions.
  - 11. The process according to claim 1, wherein said gallium-containing feedstock consists essentially of gallium-containing ntride and/or its precursors.
  - 12. The process according to claim 11, wherein said precursors are selected from the group consisting of gaillum azides, gaillum limides, gaillum amide-imides, gaillum amides, gaillum hydrides, gaillum-containing alloys, and metallic gaillum and optionally corresponding compounds of other elements of Group XIII (according to IUPAC, 1989).

- 13. The process according to claim 1, wherein sald crystallization seed has at least a crystalline layer of gallium-containing nitride.
- 14. The process according to claim 1, wherein said crystallization seed has at least a crystalline layer of gallium-containing nitride with a dislocation density below 10<sup>6</sup>/cm<sup>2</sup>.
- 15. The process according to claim 1, wherein said crystallization of a gallium-containing nitride takes place at a temperature from 100 to 800°C, preferably 300 to 600°C, more preferably 400 to 550°C.
- 16. The process according to claim 1, wherein said crystallization of a gallium-containing nitride takes place at a pressure from 100 to 10000 bar, preferably 1000 to 5500 bar, more preferably 1500 to 3000 bar.
  - 17. The process according to claim 1, wherein the content of alkali metal lons in the supercritical solvent is controlled so as to provide adequate levels of solubility of said feedstock as well as of said gallium-containing feedstock.
  - 18. The process according to claim 1, wherein the molar ratio of the moles of said alkali metal ions to the moles of the supercritical solvent is controlled within the range of 1:200 to 1:2, preferably 1:100 to 1:5, more preferably 1:20 to 1:8.
  - 19. An apparatus for obtaining of a monocystalline gallium-containing nbride crystal, comprising an autoclave 1 for producing supercritical solvent, equipped with an installation 2 or establishing a convective flow, the autoclave being mounted inside a furnace or set of furnaces 4 which are equipped with heating devices 5 and/or cooling severes 6.
- 20. The appeartus according to claim 19, wherein said furnace or set of furnaces 4 has a high-temperature zone coinciding with the crystalization zone 14 of asidutucloser a cupipped with heating devices 5, and low temperature zone coinciding with the dissolution zone 13 of the autoclave 1 equipped with heating devices 5 and/or cooling devices 6.
  - 21. The apparatus according to claim 20, wherein said furnace or set of furnaces 4 has a high-temperature zone coinciding with the crystallization zone 14 of said autoclave 1 equipped with healting devices 6 and/or cooling devices 6, as well as a low-temperature zone coinciding with the dissolution zone 13 of the autoclave 1 equipped with healting devices 6 and/or cooline devices 6.

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- 22. The apparatus according to claim 20, wherein said installation 2 is in the form of a horizontal baffle or horizontal baffles 12 having central and/or circumferential openings, separating the crystallization zone 14 from the dissolution zone 15.
  - 23. The apparatus according to claim 20, wherein feedstock 16 is placed in the autoclave 1 in the dissolution zone 13, and said a crystallization seed 17 is placed in the crystallization zone 14, and said convective flow between the zones 13 and 14 is established by said installation 2.
  - 24. The apparatus according to claim 23, wherein said dissolution zone 13 is located above said horizontal baffle or horizontal baffles 12, whereas said crystallization zone 14 is located below said horizontal baffle or horizontal baffles
- 25. A process for preparing a bulk monocrystalline gallium-containing nitride crystal in an autoclave, which comprises the steps of (i) providing a supercritical ammonia solution containing ions of alkall metal and gallium in a soluble form by introducing a gallium-containing featock to supercritical ammonia solutor containing fore of alkall metals, in which solubility of gallium-containing nitride shows a negative temperature coefficient in said supercritical ammonia solution, and (ii) crystalizing said gallium-containing nitride selectively on a crystallization seed from said supercritical ammonia solution, and (iii) crystalizing sold gallium-containing nitride selectively on a crystallization seed from said supercritical ammonia solution by means of the negative temperature coefficient of solubility.
- 26. A process for preparing a bulk monocystalline gallium-containing nitride in an autoclave, which comprises the steps of (i) providing a supercritical ammonia solution containing lons of alkall metal and gallium in a soluble form by introducing a gallium-containing feedstock into a supercritical ammonia solvent containing inos of alkall metals, in which solubility of gallium-containing influes shows a positive pressure coefficient in said supercritical ammonia solution, and (ii) crystallizing said gallium-containing ritirde selectively on a crystallization seed from said supercritical ammonia solution to means of the positive pressure coefficient of solubility.

- 27. A process for preparing a bulk monocrystalline gallium-containing nitride in an autoclave according to Claim 25 or 26, wherein said gallium-containing nitride is GaN.
- 28. A process for preparing a bulk monocrystalline gallium-containing nitride in an autoclave according to Claim 25 or 26, wherein said ion of alkali metal is selected from the group consisting of Li+, Na+, and K+.
- 29. A process for preparing a bulk monocrystalline gallium-containing nitride in an autoclave according to Claim 25 or 26, wherein said ions of alkali metals are introduced in the form of mineralizers selected from alkali metals and compounds thereof, such as azides, nitrides, amidos, amido-imides, imides, and/or hydrides, for forming an ammonobasic supercritical ammonia solution, which does not contain ions of halogens.

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- 30. A process for preparing a bulk monocrystalline gallium-containing nitride in an autoclave according to Claim 25 or 26, wherein said gallium-containing nitride is dissolved in said supercritical ammonia solvent in the form of gallium complex compounds containing alkali metals and NH<sub>2</sub> andfor its derivatives.
- 31. A process for preparing a bulk monocrystalline gaillum-containing nitride in an autoclave according to Claim 25 or 26, wherein said gailium complex compound in said supercritical ammonia solution is formed from a dissolution of GaN and/or dissolution of metallic Ga with supercritical ammonia solvent.
- 20 32. A process for preparing a supercritical ammonia solution containing gallium-containing intride, which comprises the steps of (i) providing a supercritical ammonia solvent by means of adjusting a temperature and/or pressure in an autoclave and (ii) dissolving a precursor of gallium-containing nitride in said supercritical ammonia solvent to form soluble gallium complex compounds at a temperature lower than that at which dissolving of gallium-containing nitride takes place effectively.
  - 33. A process for preparing a supercritical ammonia solution containing gallium-containing nitride according to claim 32, wherein the step of dissolving said precursor in said supercritical ammonia solvent is carried out at a temperature of 150 to 300°C.
- 34. A process for controlling the recrystallization of a gallium-containing nitride in a supeordical ammonia solution, which comprises the steps of (i) preparing a supeordical ammonia solution containing soluble gallium complex compounds formed by diseovings of gallium-containing nitride feedstock in an autoclave and (ii) decreasing the solubility of said gallium-containing nitride in the supeordical ammonia solution by increasing the temperature above that at which dissolving of the gallium-containing nitride feedstock is carried out.
  - 35. A process for controlling the recrystallization of a gallium-containing nitride in a supercritical ammonia solution which comprises the sleps of (i) preparing a supercritical ammonia solution containing soluble gallium complex compounds formed by discoving of gallium-containing nitride fleedstock in the dissolution zone and (ii) controlling over-saturation of said supercritical ammonia solution with respect to the crystallization seed, while maintaining a temperature in the crystallization zone lower than that in the dissolution zone.
  - 36. A process for controlling recrystallization of a gallium-containing nitride in a superofitical ammonia solution according toolaim 35, in which over-saturation of said supperoficial solution with respect to said crystallization seed is maintained below the level where the phenomenon of spontaneous nucleation of gallium-containing nitride appears.
  - 37. A process for controlling recrystallization of a gallium-containing nitride in a supercritical ammonia solution according to claim 35, in which over-saturation of said supercritical ammonia solution with respect to said crystallization seed, is controlled by adjusting pressure and composition of the supercritical ammonia solvent.
- 38. A process for controlling recrystallization of a gallium-containing nitride in a supercritical ammonia solution according to claim 35, in which over-saturation of said supercritical ammonia solution is controlled by adjusting the crystallization temperature.
  - 39. A process for controlling recrystallization of a gallium-containing nitride in a supercritical ammonia solution according to claim 35, in which over-saturation of said supercritical ammonia solution is controlled by adjusting the temperature difference between the dissolution zone and the crystallization zone.
    - 40. A process for controlling recrystallization of a gallium-containing nitride in a supercritical ammonia solution according

to claim 35, in which over-saturation of said supercritical ammonia solution is controlled by adjusting the rate of chemical transport.

- 41. A process for controlling recrystallization of a gallium-containing nitride in a supercritical ammonia solution according to claimi34, in which controlling sububility of said gallium-containing nitride in the supercritical ammonia solution is carried out by adulating the convective flow between the dissolution zone and the crystallization zone.
  - 42. A process for controlling recrystallization of a gallium-containing nitride in a supercritical ammonia solution according to claim 34, in which controlling solubility of said agailum-containing nitride in the supercritical ammonia solution is carried out by adjusting the opening ratio of a bailtie or bailties between the dissolution zone and the crystallization zone.
- 43. Substrate for epitaxy crystallized on the surface of a crystallization seed, especially a substrate for nitride semiconductor layers, wherein the substrate has a layer of bulk monocrystaline gallium-containing nitride, having a surface area of more than 2 cm<sup>2</sup> and having a dislocation density of less than 10<sup>2</sup> / cm<sup>2</sup>.
- Substrate for epitaxy according to claim 43, wherein the layer of bulk monocrystalline gelfium-containing nitride has
  the general formula Al<sub>2</sub>Ga<sub>1-xy</sub>In<sub>2</sub>N,
  where ®cx-1, 0Syx-1, and 0Sx+yx-1.
- 45. Substrate for epitaxy according to claim 43, wherein the substrate contains alkali elements in an amount of more than about 0.1 ppm.
- 46. Substrate for epitaxy according to claim 43, wherein the layer of bulk monocrystalline gallium-containing nitride has a halogen content that does not exceed about 0.1 ppm.
  - 47. Substrate for epitaxy according to claim 43, wherein the layer of bulk monocrystalline gallium-containing nitride has a volume of more than 0.05 cm<sup>3</sup>.
- 48. Substrate for epitaxy according to claim 43, wherein in the layer of bulk monocrystalline gallium-containing nitride has a full width at half maximum (FWHM) of X-ray rocking curve from (0002) plane of less than 600 arcsec.
  - Substrate for epitaxy according to claim 43, wherein the layer of bulk monocrystalline gallium-containing nitride
    additionally contains at least one donor depart and/or at least one acceptor depart and/or at least one magnetic
    depart in a concentration from 107 to 1021 / or m<sup>2</sup>.
  - 50. Substrate for epitaxy according to claim 43, wherein the layer of bulk monocrystalline gallium-containing nitride contains Al and/or In and the molar ratio of Ga to Al and/or In is more than 0.5.
- 51. Substrate for epitaxy according to claim 43, wherein the layer of bulk monocrystalline gallium-containing nitride is crystallized on the surface of a crystallization seed of gallium-containing nitride having a dislocation density of less than 10<sup>9</sup> / cm<sup>2</sup>.
- 52. Substrate for epitaxy according to claim 43, wherein the layer of bulk monocrystalline gallium-containing nitride has a dislocation density of less than 10<sup>4</sup> / cm<sup>2</sup> and a full width at half maximum (FWHM) of X-ray rocking curve from (0002) olane of less than 60 acresec.
  - 53. A process for obtaining a gallium-containing nitride crystal, comprising the steps of:
  - (i) providing a gallium-containing feedstock, an alkali metal-containing component, at least one crystallization seed and a nitrogen-containing solvent in at least one container;
    - (ii) bringing the nitrogen-containing solvent into a supercritical state:
    - (iii) at least partially dissolving the gallium-containing feedstock at a first temperature and at a first pressure; and
    - (iv) crystallizing gallium-containing nitride on the crystallization seed at a second temperature and at a second pressure while the nitrogen-containing solvent is in the supercritical state;

wherein at least one of the following criteria is fulfilled:

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- (a) the second temperature is higher than the first temperature; and
- (b) the second pressure is lower than the first pressure.

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- 54. The process according to claim 53, wherein the at least one container is an autoclave.
- 55. The process according to claim 53, wherein the gallium-containing feedstock is at least partially dissolved before step (iv).
- 56. The process according to claim 53, wherein the gallium-containing feedstock is at least partially dissolved during step (iv).
- 57. The process according to claim 53, wherein the process is conducted in a container having a dissolution zone at the first temperature and a crystallization zone at the second temperature and wherein the second temperature is higher than the first temperature.
- 58. The process according to claim 57, wherein the difference in temperature between the dissolution zone and crystallization zone is selected so as to ensure convective transport in the supercritical solution.
- 59. The process according to claim 58, wherein the difference in temperature between the second temperature and the first temperature is at least 1 °C.
  - The process according to claim 59, wherein the difference in temperature between the second temperature and the first temperature is from about 5 to about 150 °C.
- 61. The process according to claim 53, wherein the gallium-containing nitride has the general formula Al<sub>x</sub>Ga<sub>1-x-y</sub>in<sub>y</sub>N,
  where 0≲x<1, 0≲x<1, and 0≤x+y<1.</li>
- The process according to claim 61, wherein the gallium-containing nitride has the general formula Al<sub>x</sub>Ga<sub>1-x-y</sub>in<sub>y</sub>N, where 0≤x<0.5 and 0≤y<0.5.</li>
- 63. The process according to claim 61, wherein the gallium-containing nitride is gallium nitride.
  - 64. The process according to claim 53, wherein the gallium-containing nitride further contains at least one donor dopant, at least one acceptor dopant, at least one magnetic dopant or mixtures thereof.
  - 65. The process according to claim 53, wherein the gallium-containing feedstock comprises at least one compound selected from the group consisting of gallium nitride, gallium azides, gallium irrides, gallium aniido-irrides, gallium hydrides. asilum-containing alloys, metallic gallium and mixtures thereof.
- 66. The process according to claim 65, wherein the gallium-containing feedstock comprises metallic gallium and gallium nitride.
  - 67. The process according to claim 65, wherein the feedstock further comprises an aluminium feedstock, an indium feedstock or mixtures thereof, wherein the feedstocks are selected from the group consisting of nitrides, azides, imides, emid-imides, privides, alloys, metallic aluminium, and metallic indium.
    - 68. The process according to claim 53, wherein the alkali metal-containing component is at least one metallic alkali metal or at least one alkali metal salt.
- 69 69. The process according to claim 68, wherein the alkali metal in the alkali metal-containing component is lithium, sodium, potassium or cesium.
  - 70. The process according to claim 69, wherein the alkali metal in the alkali metal-containing component is sodium or notes six in.
  - 71. The process according to claim 68, wherein the alkali metal salt is an amide, an imide or an azide.
  - 72. The process according to claim53, wherein a surface of the at least one crystallization seed is a crystalline layer of

a gallium-containing nitride.

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- The process according to claim 72, wherein the gallium-containing nitride of the crystalline layer has the general formula Al<sub>x</sub>Ga<sub>1-x-y</sub>In<sub>y</sub>N, where 0≤x<1, 0≤y<1, and 0≤x+y<1.</li>
  - 74. The process according to claim 72, wherein the crystalline layer has a dislocation density of less than 10<sup>6</sup> / cm<sup>2</sup>.
  - 75. The process according to claim 53, wherein the nitrogen-containing solvent is ammonia, a derivative thereof, or mixtures thereof.
- 76. The process according to claim 53, wherein the first temperature and the second temperature are from about 100 °C to about 800 °C and wherein the second temperature is at least 1 °C higher than the first temperature.
- 77. The process according to claim 76, wherein the first temperature and the second temperature are from about 300 °C to about 600 °C.
  - 78. The process according to claim 77, wherein the first temperature and the second temperature are from about 400 °C to about 550 °C.
- 79. The process according to claim 53, wherein the first pressure and the second pressure are the same and are from about 1000 bar (105 kPa) to about 10 000 bar (106 kPa).
  - 80. The' process according to claim 79, wherein the first pressure and the second pressure are the same and are from about 1000 bar (105 kPa) to about 5500 bar (5.5x105 kPa).
  - 81. The process according to claim 80, wherein the first pressure and the second pressure are the same and are from about 1500 bar (1.5x10<sup>5</sup> kPa) to about 3000 bar (3x10<sup>5</sup> kPa).
  - The process according to claim 53, wherein step (iv) is conducted so that the crystallization selectively takes place
    on the crystallization seed.
    - 83. A process for preparing a gallium-containing nitride crystal comprising the steps of:
    - (i) providing a gallium-containing feedstock comprising at least two different components, an alkali metal-containing component, at least one crystallization seed and a nitrogen-containing solvent in a container having a dissolution zone and a crystallization zone, whereby the gallium-containing feedstock is provided in the dissolution zone and the at least one crystallization seed is provided in the crystallization zone;
      - (ii) subsequently bringing the nitrogen-containing solvent into a supercritical state;
- (iii) subsequently partially dissolving the gallium-containing feedstock at a dissolution temperature and at a dissolution pressure in the dissolution zone, whereby a first component of the gallium-containing feedstock is substantially completely dissolved and a second component of the gallium-containing feedstock as well as the crystallization seed(s) remain substantially undissolved so that an undersaturated or saturated solution with respect to allium-orbatining nitride is obtained;
- (iv) subsequently setting the conditions in the orgatalization zone at a second temperature and at a second pressure so that over-asturation with respect to gallium-containing nitride is obtained and orgatalization of gallium-containing nitride occurs on the at least one crystalization seed and setting the conditions in the dissolution zone at a first temperature and at a first pressure so that the second component of the gallium-containing feedstock is dissolved;
- 50 wherein the second temperature is higher than the first temperature.
  - 84. The process according to claim 83, wherein the first component of the gallium-containing feedstock is metallic gallium and the second component of the gallium-containing feedstock is gallium nitride.
- 85. The process according to claim 83, wherein the crystallization is conducted so that it selectively takes place on the crystallization seed.
  - 86. The process according to claim 83, wherein the first temperature and the first pressure in the dissolution zone and

the second temperature and the second pressure in the crystallization zone are selected so that the concentration of gallium in the over-saturated solution remains substantially the same during crystallization.

- 87. The process according to claim 83, wherein the container comprises at least one baffle between the dissolution zone and the crystallization zone.
  - 88. The process according to claim 87, wherein the at least one baffle has a central opening, circumferential openings or a combination thereof.
- 89. A gallium-containing nitride crystal obtainable by a process according to any one of claims 53 to 88.

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- A gallium-containing nitride crystal having a surface area of more than 2 cm<sup>2</sup> and having a dislocation density of less than 10<sup>6</sup> / cm<sup>2</sup>.
- 91. A gallium-containing nitride crystal having a thickness of at least 200 μm and a full width at haif maximum (FWHM)
  of X-ray rocking curve from (0002) plane of 50 arcsec or less.
  - 92. The gallium-containing nitride crystal according to claim 91 wherein the thickness is at least 500 μm.
- 33. The gallium-containing nitride crystal according to any of claims 89 to 92, wherein the gallium-containing nitride crystal has the general formula Al<sub>x</sub>Ca<sub>1-xy</sub>In<sub>x</sub>N, where 0≤x<1, 0≤y<1, and 0≤x+y<1.</li>
  - 94. The gallium-containing nitride crystal according to any of claims 89 to 92, wherein the gallium-containing nitride crystal contains alkali elements in an amount of more than about 0.1 ppm.
  - 95. The gallium-containing nitride crystal according to any of claims 89 to 92, wherein the gallium-containing nitride crystal has a halogen content of about 0.1 ppm or less.
  - 96. The gallium-containing nitride crystal according to any of claims 89 to 92, wherein the gallium-containing nitride crystal has a volume of more than 0.05 cm<sup>3</sup>.
    - 97. The gallium-containing nitride crystal according to any of claims 89 to 92, wherein the gallium-containing nitride crystal contains at least one element selected from the group consisting of Ti, Fe, Co, Cr, and Ni.
- 5 99. The gallium-containing nitride crystal according to any of claims 89 to 92, wherein the gallium-containing nitride crystal additionally contains at least one donor dopart and/or at least one acceptor dopart and/or at least one magnetic dopart in a concentration from 1017 to 1021 (cm<sup>3</sup>).
- 99. The gallium-containing nitride crystal according to any of claims 89 to 92, wherein the layer of gallium-containing of nitride crystal further contains A1 and/or In and the molar ratio of Ga to Al and/or In is more than 0.5.
  - 100.The gallium-containing nitride crystal according to any of claims 89 to 92, wherein the gallium-containing nitride crystal contains a seed.
- 45 101.The gallium-containing nitride crystal according to any of claims 89 to 92, wherein the gallium-containing nitride crystal is monocrystalline.
- 102.An apparatus for obtaining a gailium-containing nitride crystal comprising an autoclave 1 having an internal space and comprising at least one device 4, 5 for heating the autoclave be at least two conse having different therepartures, or wherein the autoclave comprises a device which separates the internal space into a dissolution zone 13 and a crystallization zone 14.
  - 103. The apparatus according to claim 102, wherein the at least one device is for heating the autodave to two zones having different temperatures and the two zones coincide with the dissolution zone 13 and the crystallization zone 14.
  - 104. The apparatus according to claim 102, wherein the device which separates the internal space is at least one baffle 12 having at least one opening.

- 105.The apparatus according to claim 104, wherein the at least one baffle 12 has a central opening, circumferential openings or a combination thereof.
- 106.The apparatus according to claim 102, wherein the crystallization zone 14 is provided with a heating device 5 for heating the crystallization zone 14 to a temperature higher than the temperature of the dissolution zone 13.
- 107.The apparatus according to claim 102, wherein a seed-holder 18 is provided in the crystallization zone 14 and a feedstock-holder 19 is provided in the dissolution zone 13.
- 10 108. The apparatus according to claim 104, wherein the baffle(s) is/are in a horizontal position and wherein the dissolution zone 13 is located above said horizontal baffle or horizontal baffles 12, whereas said crystallization zone 14 is located below said horizontal baffle or horizontal baffles 12.

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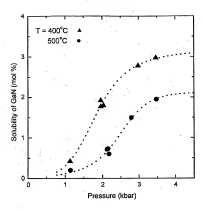
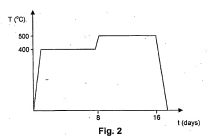


Fig. 1



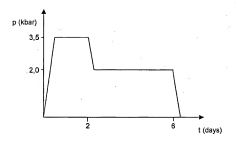


Fig. 3

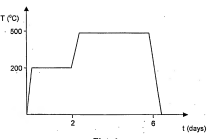


Fig. 4

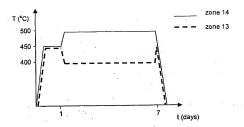
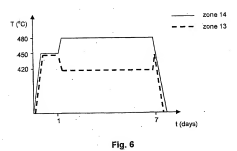


Fig. 5



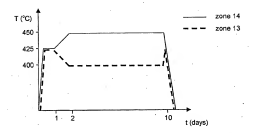


Fig. 7

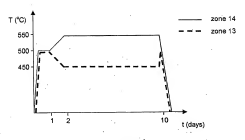
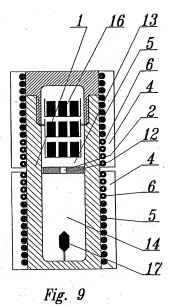
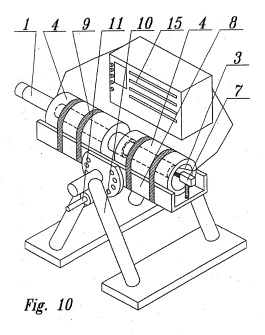


Fig. 8





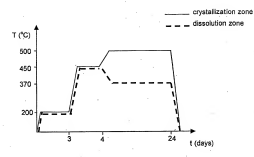


Fig. 11

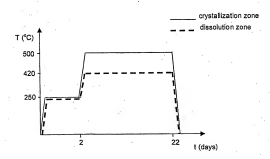


Fig. 12

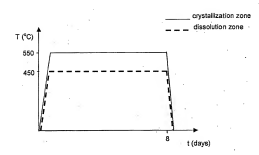


Fig. 13

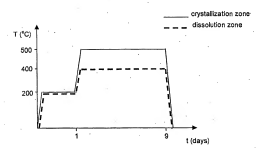


Fig. 14

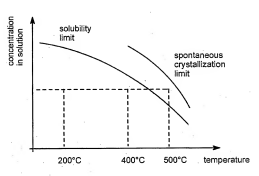


Fig. 15

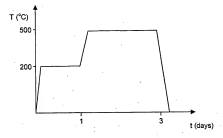


Fig. 16

# REFERENCES CITED IN THE DESCRIPTION

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